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RECORDING MEDIUM

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[There are no amendments to this patent.]

Claim

A recording medium in which a monomolecular film or a built-in monomolecular film layer comprising a host molecule having a hydrophilic group, a hydrophobic group, and an inclusion group, and a guest molecule having photodimeric properties and is enclosed in the above-mentioned host molecule and is produced on a carrier to form a recording layer, and recording is performed by means of light.

Applicant:

Agent:

Detailed explanation of the invention

(1) Field of technology

The present invention pertains to a recording medium in which recording is performed by utilizing the chemical or physical change in a monomolecular film consisting of an inclusion complex or built-in monomolecular film layer.

(2) Background of the technology

A variety of recording mediums having an organic compound as the recording element are known.

For example, an optical recording medium in which a thin-film organic compound is used as the recording layer is disclosed in, for example, Japanese Kokai Patent Application Nos. Sho 56[1981]-16948 and Sho 58[1983]-125246. Each of the above-mentioned cases pertains to a laser recording medium in which an organic colorant is used as the recording medium, and recording and playback are achieved with a laser beam. In particular, the medium disclosed in Japanese Kokai Patent Application No. Sho 58[1983]-125246 has a recording layer consisting of a thin film of a cyanine dye indicated by general formula (I) below.

General formula (I)

The cyanine dye indicated by chemical structure (I) is coated onto a plastic base to form a thickness of 1000 Å or less, for example, a thickness of approximately 300 Å, by means of a rotary coating machine, and a thin film is produced. When the distribution of the molecular orientation inside the film is random, when photoradiation is applied, optical scattering occurs within the film, and the degree of chemical reaction that occurs upon application of each type of radiation varies on a microscopic level. For this reason, a recording medium with a uniform molecular distribution and orientation within the molecular layer is desirable, and a reduced film thickness is required for increased recording density. However, when a coating method is used, the limitation in thickness of the film is approximately 300 Å, and random molecular distribution and orientation within the film was difficult to achieve.

In Japanese Kokai Patent Application Nos. Sho 56[1981]-42229 and Sho 56[1981]-43220, a built-in film of diacetylene compound with a significant photonic effect and excellent resolution is suggested as a resist material that can be used in thin-film electrooptical devices, electroacoustic devices, piezopyroelectric devices, etc., as well as being used in resist materials.

Recently, an improvement in the method of manufacturing diacetylene compound built-in film has been disclosed in Japanese Kokai Patent Application No. Sho 58[1983]-111029. The diacetylene compound built-in film produced on a base by the method described in the above-mentioned invention undergoes polymerization upon being exposed to ultraviolet [radiation] and produces a diacetylene compound polymer film, or masking can be

used and the ultraviolet [radiation] applied to produce localized polymerization and the nonpolymerized portions are removed to produce a pattern, and the result is used as a thin-film optical device or integrated circuit element.

However, all of the above-mentioned methods are limited to diacetylene compounds, and the possibility of erasing the recording when used as a thin-film optical device is not discussed at all.

Meanwhile, as a means to overcome the above-mentioned limitations, an optical recording medium that can be used repeatedly has been produced by forming a recording layer consisting of a monomolecular film or monomolecular built-in film comprised of at least a photopolymeric monomer containing a hydrophilic [group], a hydrophobic [group], and at least one unsaturated bond in the molecule on a base and is disclosed in Japanese Patent Application No. Sho 58[1983]-190932.

In all of the above-mentioned diacetylene compound built-in films and monomolecular films or monomolecular built-in films comprised of a photopolymeric monomer, a method of manufacturing in which a hydrophilic group and hydrophobic group are introduced to a photoreactive compound, which is deposited directly on a base, is used. As a result, it is difficult to produce films with different functions using a simple method, and a reduction in the photoreactivity of the film results from introduction of the hydrophilic group and hydrophobic group. Furthermore, to control the molecular orientation inside the film, which is very important for high-density recording, a very complex operation is required.

In an effort to overcome the above-mentioned limitations in the conventional technology and as a result of their research on 1) a method of producing a variety of functional films using a relatively simple method, 2) a method in which the above-mentioned film formation is carried out in such a manner that the functional properties of the molecule are not diminished even when a thin film is used, and 3) a method in which orientation of the film molecular structure in the in-plane direction of the film is carried out to provide a high orientation without a special operation; thereby, the present invention was accomplished. Furthermore, it was easily possible to produce a high-quality recording medium with high sensitivity and high resolution using the above-mentioned method.

(3) Disclosure of the invention

The objective of the present invention is to produce a high-density recording medium capable of chemical change or physical change on the molecule level as a result of exposure to light.

Also, the objective is to produce a superior medium compared to those produced by conventional methods with regard to the molecular orientation within the plane of the medium which is an important factor for high-density recording on the molecular level. Furthermore, the objective is to produce a medium with a variety of properties through a relatively simple change in the operation during production of the above-mentioned recording medium.

The above-mentioned objectives of the present invention can be achieved by the present invention described below.

A recording medium in which a monomolecular film or a built-in monomolecular film layer comprising a molecule (host molecule) having a hydrophilic group, a hydrophobic group, and an inclusive group (inclusion group), and a molecule having photodimeric properties (guest molecule) consisting of a different material which is enclosed in the above-mentioned host molecule is produced on a carrier to form a recording layer, and recording is performed by means of light.

The material that comprises the recording layer of the present invention consists of two types of molecules, a molecule having at least one group containing a hydrophilic group, a hydrophobic group, and a group whereby inclusion with a different molecule is made possible (which is referred to as the host molecule), and a different type of molecule is enclosed in the above-mentioned host molecule (which is referred to as the guest molecule). When the monomolecular film, or built-in monomolecular film layer of the inclusion complex consisting of the above-mentioned host molecule and guest molecule is deposited on a carrier, the recording medium of the present invention can be produced. Between the above-mentioned two types of molecules, it is necessary for the guest molecule to be able to achieve dimerization upon exposure to light.

Host molecules that can be used in the present invention, as described above, are molecules having a hydrophilic group, a hydrophobic group, and a group capable of forming at least one inclusion complex with a different type of molecule, and it is not especially limited. As typical examples of structural

elements capable of forming the hydrophilic group or hydrophobic group within the molecule, a variety of commonly known hydrophilic groups and hydrophobic groups can be mentioned. The group capable of forming an inclusion complex with a different type of molecule can be produced by introduction of hydroxy groups, carbonyl groups, carboxyl groups, ester groups, amino groups, nitrile groups, thiol groups, imino groups, etc. In the following, the above-mentioned host molecule is explained in specific terms using host molecules having hydroxy groups indicated by general formulas (IIa)-(IIc) below as examples.

(In this case, X = H or C_6H_5 .)

In other words, having a hydrophilic group and hydrophobic group within the molecule means, for example, in the above-mentioned chemical structure, a hydrophilic group exists on either R_1 or R_2 , and a hydrophobic group exists on the other, or both the R_1 group and the R_2 group exhibit hydrophilic properties or hydrophobic properties in relation to groups other than those two. Regarding the structure of R_1 and R_2 , when a hydrophobic group is to be introduced, a long-chain alkyl group with 5-30

carbon atoms is especially suitable, and when a hydrophilic group is to be introduced, a fatty acid with 1-30 carbon atoms is especially suitable.

For specific examples of the host molecule used in the present invention, acetylene diol derivatives (Nos. 1-6 and Nos. 16-21), diacetylene diol derivatives (Nos. 7-12 and Nos. 22-27), hydroquinone derivatives (Nos. 13-15, and Nos. 28-30), etc., can be mentioned as suitable examples. It should be noted that in the following, m and n each represent a positive number, Z represents $-CH_3$ or -COOH, and Ph represents $-C_6H_5$.

[Examples of acetylene diol derivatives]

No. 1

$$CH_{3}-(CH_{2})_{20}-C-C \equiv C-C-(CH_{2})_{20}-COOH$$

OH
OH
OH

No. 2

$$CH_{3}-(CH_{2})_{22}-C-C=C-\frac{1}{C}-(CH_{2})_{22}-COOH$$
OH
OH
OH
OH

 $30 \ge m + n \ge 11$. $n \ge 0$

No. 3

$$CH_{s}-(CH_{t})_{2s}-O-\bigcirc - CO-C - C-C-C-\bigcirc OH$$

30 ≥ m + n ≥ 8, n ≥ 1

. خ. 5

$$CH_{s}-(CH_{t})_{m} \xrightarrow{O} -O - \overset{!}{C} - C = C - \overset{!}{C} - \overset{!}{O} - (CH_{t})_{a} - COOH$$

$$OH \qquad OH$$

$$30 \ge m + n \ge 8. \quad n \ge 0$$

No. 6

$$CH_3 - (CH_1)_m - \bigcirc -O - \stackrel{Pb}{C} - C = C - \stackrel{C}{C} - \bigcirc -(CH_1)_a - COOH$$

$$OH \qquad OH$$

$$30 \ge m + n \ge 8, \quad n \ge 0$$

[Examples of diacetylene diol derivatives]

. io. 7

$$H H H H CH3-(CH3)70-C-C = C-C = C-C-(CH3)0-COOH OH$$

No. 8

No. 9

No. 10

$$CH_3 - (CH_1)_m - O - \bigcirc - \stackrel{Pb}{\bigcirc} - \stackrel{Pb}{\bigcirc} - C - C = C - C = C - C - \bigcirc - O - (CH_1)_m - COOH$$

$$0H \qquad OH$$

$$30 \ge m + n \ge 5, \quad n \ge 1$$

No. 12

 $30 \ge m + n \ge 5$, $n \ge 0$

[Examples of hydroquinone derivatives]

No. 13

 $30 \ge m + n \ge 13$, $n \ge 0$

No. 14

 $30 \ge m + n \ge 9$, $n \ge 1$

No. 15

30 ≥ m + m ≥ 9, m ≥ 0

[Examples of acetylene diol derivatives]

No. 16

$$Z - (CH_1)_a - \overset{i}{C} - C = C - \overset{i}{C} - (CH_1)_a - Z$$

30 ≥ a ≥ 5

No. 17

$$Z - (CH_1)_a - \overset{Pb}{\overset{!}{C}} - C \equiv C - \overset{!}{\overset{!}{C}} - (CH_1)_a - Z$$

30 ≥ a ≥ 5

No. 19

$$Z - (CH_1)_a - O - \bigcirc \bigcirc - \stackrel{Pb}{\stackrel{c}{\stackrel{c}{\circ}}} - C = C - \stackrel{c}{\stackrel{c}{\stackrel{c}{\circ}}} - \bigcirc - O - (CH_1)_a - Z$$

$$\stackrel{30}{\stackrel{>}{\circ}} = 2 4$$

No. 20

No. 21

30 ≥ a ≥ 4

With the exception of substitution of a long-chain alkyl group or long-chain carboxylic acid for the host molecule, and introducing hydrophilic properties or hydrophobic properties, the above-mentioned compounds themselves are known compounds; furthermore, formation of a crystalline inclusion complex of the host molecule that is not modified with a long-chain alkyl group, etc., is also described in the Journal of the Chemical Society of Japan, No. 2, pp. 239-242 (1983).

For the guest molecule capable of producing an inclusion complex with the above-mentioned host molecules, in general, molecules capable of forming a strong hydrogen bond with the host molecule are desirable. For this reason, as described above, when the host molecule has a hydroxy group as the inclusion group, aldehyde, ketone, amine, sulfoxide, etc., can be mentioned as guest molecules. Furthermore, for guest molecules, different types of halogen compounds, or π -electron compounds, in other words, alkene, alkyne, arene, etc., can be selected. In either case, a molecule having a structure that makes it possible for the inclusion complex produced to have a desired image formation ability [is selected].

In the following, specific examples of guest molecules that can be used for a recording medium that utilizes a photodimerization reaction of the guest molecule, olefin compounds (Nos. 31-34), diolefin compounds (Nos. 35-38), anthracine derivatives (No. 39), 2-aminopyridinium (No. 41), etc., can be mentioned.

[Examples of olefin compounds]*

^{*[}Editor;s note: compound Nos. 22-30 are not included in the original.]

No. 33

No. 34

$$R_1OOC - CH = CH - COOR_1$$

 $(R_1 = R_2 = CH_3 \pm f_2 + R_1 = CH_3, R_2 = H_3)$

Key: 1 Or

[Examples of diolefin compounds]

No. 35

Key: 1 For

№. 36

Key: 1 For

No. 37

CN

$$R - CH = CH - COCH_{1} - COOCH_{2}$$

No. 38

[Examples of anthracene derivatives]

[Examples of acrydinium derivatives]

[2-aminopyridinium]

As a method of producing the monomolecular film or built-in monomolecular film layer of the inclusion complex comprised of above-mentioned host molecules and guest molecules, for example, the Langmuir-Blodgett method (LB method) developed by I. Langmuir et al. can be used. The LB method is a method in which a monomolecular film or built-in film of the monomolecular layer is produced by utilizing the tendency of molecules to form a monomolecular layer on the surface of water with the hydrophilic

group facing downward when an appropriate balance is retained in a molecular structure having a hydrophilic group and hydrophobic group in the molecule (that is, balance of the two properties). The monomolecular layer on the surface of the water is a two-dimensional system. When the molecules are distributed at random, the equation of a two-dimensional ideal gas can be established between surface [area] A and the surface pressure π per molecule,

$\pi A = kT$

and forms a "gaseous film". In this case, k is Boltzmann's constant and T is the absolute temperature. When A is adequately reduced, the intramolecular interaction increases and forms a "condensed film" (or solid film) consisting of a two-dimensional solid material. The condensed film can be transferred to the surface of a carrier comprised of a different material or having a different shape such as a glass base, one layer at a time. As a specific example of the method of producing the monomolecular film (which is referred to as a single complex molecular film), or single complex molecular built-in film of the host molecule that includes the guest molecule of the present invention, five methods A-E shown below can be mentioned.

(A) The host molecule and the guest molecule of the target inclusion complex are dissolved in a solvent. [The complex] is then formed in an aqueous phase and the inclusion complex is precipitated in the form of a film. In this case, when the structure of the host molecule includes a hydrophilic group (carboxyl group) and a hydrophobic group (alkyl group) on both ends of the molecule as indicated by compound Nos. 1-15, the

inclusion complex precipitated on the aqueous phase is developed on the aqueous phase with the hydrophilic group of the host molecule facing the aqueous phase regardless of the hydrophilic and hydrophobic properties of the guest molecule. On the other hand, when the host molecule takes on the structure indicated by compound Nos. 16-30, $Z=-CH_3$, wherein both ends of the molecule consist of only hydrophobic groups, and the inclusion complex develops on the aqueous phase with the hydrophilic group of the guest molecule facing the aqueous phase as indicated by Figure 4. Also, for Z=-COOH, wherein both ends of the molecule consist of hydrophilic groups alone, the inclusion complex develops on the aqueous phase with the hydrophilic group of the host molecule facing the aqueous phase, as indicated by Figure 5.

Subsequently, a partition board (or a float) is provided so that spreading of the precipitate on the aqueous phase does not occur freely, the area of spread is restricted, the state of spreading of the film material is controlled, and the surface pressure π in proportion to the state of spreading is obtained. The partition board is then moved, the area of spread is reduced, the state of spread in the film material is controlled, the surface pressure is slowly increased, and a surface pressure π suitable for production of the built-in film can be obtained. The surface pressure is retained, the cleaned carrier is carefully placed in a vertical motion, and the single complex molecular film is transferred to the carrier. The single complex molecular film is produced as described above, and the above-mentioned operation is repeated. A single complex molecular

built-in film layer with a desired degree of buildup can be produced.

In order to transfer the monomolecular layer onto the surface of the carrier, in addition to the above-mentioned dipping method, a horizontal adsorption method, rotational cylinder method, etc., can be mentioned. The horizontal adsorption method is a method in which the carrier is brought into contact with the surface of the water horizontally and transfer is performed. The rotational cylinder method is a method in which a cylindrical carrier is rotated on the surface of the water and the monomolecular layer is transferred to the surface of the carrier. In the above-mentioned vertical dipping method, a monomolecular layer with the hydrophilic group of the host molecule facing the carrier side can be produced when a carrier with a surface having hydrophilic properties is pulled out of the water in a direction that crosses the surface of the When the carrier is transported in the vertical direction as described above, a single complex monomolecular layer is built up for each process. The position of the film molecules produced is reversed in the removal process and dipping process; therefore, when the above-mentioned method is used, a Y-type film, in which the hydrophobic group and hydrophobic group of the host molecule and hydrophobic group and hydrophobic group of the host molecule face each other between the layers can be produced. On the other hand, in the horizontal adsorption method, the carrier is brought into contact with the surface of the water, and transfer is performed, and a single complex molecular layer with the hydrophobic group of the host molecule facing the carrier side can be produced on the carrier. In said method, a

change in the direction of the film molecules produced does not occur even when built up, and an X-type film, in which the hydrophobic group faces the carrier in all layers, can be produced. A built-in film in which the hydrophilic is facing the carrier side in all layers is referred to as a Z-type film.

In the rotational cylinder method, a cylindrical carrier is rotated on the surface of the water, the monomolecular layer is transferred to the surface of the carrier. Transferring the monomolecular layer onto the carrier is not limited to the above-mentioned methods, and when a carrier with a large surface area is used, a method in which the carrier is extruded into the aqueous phase from a carrier roll, etc., can be used as well. Also, the orientation of the hydrophilic group or hydrophobic group facing the carrier described above is a rule, a surface treatment can be provided for the carrier, and the direction can be changed.

In the above-mentioned film formation process, control of the orientation in the in-plane direction of the film material has been mainly controlled through adjustment of the surface pressure in the past, but it has been very difficult to achieve a highly ordered orientation unless the film material is a compound with a very simple structure, for example, a straight-chain fatty acid, etc. However, in the present invention, an inclusion complex is used as a film material; thus, a film with a highly ordered orientation can be produced relatively easily. In other words, configuration between the host molecule-guest molecule, host molecule-host molecule, and guest molecule-guest molecule are defined at the point where the inclusion complex is precipitated on the aqueous phase based on the hydrogen bonds,

van der Waals forces, etc., and the host molecule and guest molecule are arranged with a crystal lattice order. Also, when the guest molecule alone has functionality, chemical modification for the guest molecule, in other words, introduction of a hydrophobic group or hydrophilic group is not performed; thus, reduction in the functionality accompanied by film formation does not occur.

- (B) A guest molecule that exhibits solubility is dissolved in the aqueous phase. Subsequently, a host molecule is dissolved with a solvent and spread onto the aqueous phase. In this case, formation of the inclusion complex takes place among the host molecule-guest molecule at the same time, and formation of a film takes place. The combination of the host molecule and guest molecule and the film formation method that follows are based on the methods described in (A).
- (C) A guest molecule that exhibits solubility is dissolved in the aqueous phase. Subsequently, the host molecules and guest molecules of the target inclusion complex are dissolved in a solvent and spread onto the aqueous phase. The combination of host molecule and guest molecule and the film formation method that follows are based on the method described in (A).
- (D) A host molecule is dissolved in a solvent and spread onto the aqueous phase. Subsequently, a closed-system device is used and the side of the gaseous phase, that is, the space inside the device, is formed into a gas atmosphere of the guest molecule. In this case, the guest molecule on the gaseous phase side is enclosed at the same time, and an inclusion complex is precipitated in a form of a film. The above-mentioned method is especially effective for compounds in which the guest molecule

has a low boiling point and is likely to vaporize, for example, acetone, etc. The combination of the host molecule and guest molecule and the film formation method that follows are based on the method described in (A).

(E) A closed-system device is used and the side of the gaseous phase, that is, the space inside the device, is filled with a gaseous atmosphere comprised of the guest molecule. Then, the host molecule and guest molecule of the target inclusion complex are dissolved in a solvent. It is then sprayed onto the aqueous phase, and an inclusion complex is precipitated in the form of a film. The combination of the host molecule and guest molecule and the film formation method that follows are based on the method described in (A).

The single complex molecular film and single complex molecular built-in film produced on the carrier using the above-mentioned methods have a high density and highly ordered orientation, and when a recording layer is made with the above-mentioned films, a recording medium having good characteristics as a high-density high-resolution recording medium that can be used for optical recording, thermal recording, electrical recording, magnetic recording, etc., can be produced.

When the single complex molecular film and single complex molecular built-in film produced above are used as the recording layer of a recording medium, recording can be performed as described below.

For example, when one of Nos. 7-15, Nos. 22-30 is used for the host molecule, and a guest molecule having a photodimerizable double bond such as those indicated by Nos. 31-38 are used in combination, an inclusion complex having a host molecule and

guest molecule composition ratio (mole ratio) of 1:2 can be produced. When the energy required to achieve polymerization such as gamma rays, X-rays, or ultraviolet [radiation] is applied to the above-mentioned single complex molecular film or single complex molecular layer built-in film according to a specified pattern, dimerization takes place between the guest molecules in the irradiated areas, as indicated by equation III shown below.

The above-mentioned reaction can occur when the distance between adjoining unsaturated bonds is 4 Å or less, but in a single complex molecular film or single complex molecular layer built-in film produced by the above-mentioned methods, not only can the dimerization product be produced easily, but also, only one type of isomer or structure among a variety of materials that are likely to be produced at the time of the dimerization reaction can be produced. In other words, the configuration between the guest molecules in the inclusion complex layer is highly ordered. Also, depolymerization does not occur after the dimerization reaction, even in the dark and areas that are not exposed to radiation remain in the monomer form; thus, a recording based on a specified pattern can be produced as indicated by Figure 1.

Reading of the recorded information can be done by, for example, irradiation or visible rays. In other words, the conjugated system of the monomer undergoes decomposition when polymerization is performed; thus, changes occur in the

absorption wavelength of the visible light. The maximum absorption wavelength shifts toward the lower wavelengths; thus, playback of the information can be achieved by reading the change in spectral absorption (Figure 2).

In addition to readout of the spectral absorption by visible light, playback can be achieved by reading the change in volume and after dimerization using the schlieren method. The above-mentioned method is especially effective when a single complex molecular film or single complex molecular built-in film of a compound has a structure that exhibits a significant difference in volume at the time of polymerization and after the dimerization. Also, instead of directly forming the single complex molecular film or single complex molecular layer built-in film on the base, an optically conductive layer such as Se, ZnO, or CdS can be produced on the base, and a single complex molecular film or single complex molecular built-in film also is produced; thus, the difference in absorption between the monomer and the dimer can be read out electronically.

The dimer has absorption at a wavelength of 270 nm based on the cyclobutane ring, and when ultraviolet [radiation] with a wavelength of 270 nm is applied, the dimer returns to the initial two monomers. Thus, erasure of the recorded information is possible (Figure 3).

Also, as the guest molecule of the photodimer, compounds such as the anthracene derivatives indicated by No. 39 can be used. In this case, the photodimerization reaction progresses according to reaction equation IV.

Also, in the case when No. 7 is used as the host molecule, polymerization takes place between host molecules in the irradiated areas as indicated by equation V when the energy required for polymerization such as X-rays, gamma rays, and ultraviolet rays is applied, and polydiacetylene is produced.

Therefore, a significant increase in the adhesion with the base is made possible when a total exposure is performed for the single complex molecular film or single complex molecular built-in film. In particular, a significant increase in the chemical resistance (solvent resistance) can be observed. When the above-mentioned total exposure is performed, when the guest molecule has photodimeric properties, dimerization also takes place in the guest molecule, but when the material is used as an

optical recording medium, as a case opposite to the example described above, when depolymerization is performed through application of light with a wavelength at the absorption wavelength of the cyclobutane ring (ultraviolet) according to a specified pattern, recording or display can be achieved.

For the above-mentioned optical recording medium, one with a film thickness of 100-3000~Å is especially desirable.

As indicated by the principle, the method of producing the film is very simple, and it is possible to produce a recording medium with the above-mentioned superior properties at low cost.

The carrier used for production of the above-mentioned single complex molecular film or single complex molecular built-in film used in the present invention is not especially limited, but when a surfactant is adsorbed on the surface of the carrier, the single complex molecular film is disturbed when the single complex molecular film layer is transferred from the surface of water, and a high-quality single complex molecular film or single complex molecular layer built-in film cannot be produced; thus, a carrier with a clean surface is used. For examples of carriers that can be used in this case, glass, metals such as aluminum, plastics, ceramics, etc., can be mentioned.

The single complex molecular film or single complex molecular built-in film formed on the carrier is firmly bonded, and peeling or delamination from the carrier hardly occurs, but in order to further increase the adhesion, an adhesive layer can be formed between the single complex molecular film or single complex molecular built-in film and the carrier. Also, the adhesion can be increased through selection of the conditions of formation of the single complex molecular film layer, for

example, concentration of the hydrogen ion in the aqueous phase, type of ion used, water temperature, carrier transport speed, etc.

It is desirable to provide a protective film on the single complex molecular film or single complex molecular built-in film from the standpoint of chemical stability of the single complex molecular film or single complex molecular built-in film, but depending on the type of film-forming molecule, a protective film is not a requirement.

In the following, the present invention is further explained in specific terms with application examples. Compound Nos. 49-54 are listed in Table I.

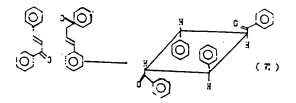
Application Example 1

An optical recording medium that utilizes the photodimerization reaction of the guest molecule (1)

Diacetylene diol indicated by No. 49 was used as a host molecule, and chalcone was used as the guest molecule. The two were dissolved in chloroform at a mole ratio of 1:2 and spread over an aqueous phase of cadmium chloride with a pH of 6.5 and concentration of 4 x 10⁴M. After removing the chloroform solvent by evaporation, the surface pressure was increased to 35 dynes/cm, and precipitation of the inclusion complex was performed in the form of a film. Subsequently, the surface pressure was kept constant, a glass board with a thoroughly clean surface having hydrophilic properties was gently moved in the

vertical direction at a rate of 7 cm/min in such a manner that the board crossed the surface of the water, and a single complex molecular film was transferred to the board. A single complex molecular film, an optical recording medium having a recording layer comprised of a single complex molecular film, and a single complex molecular built-in film formed by buildup of 3, 5, 9, 15, and 19 layers were produced. In the above-mentioned buildup process, the board was left standing for 30 min each time after being removed from the aqueous phase, and the moisture adsorbed on the board was removed by evaporation. As for the film formation machine, a Langmuir-Trough 4 produced by Joyce Corp. of England was used.

X-ray application was done for the optical recording medium produced according to a pattern, the dimerization reaction of the guest molecule indicated by equation VI was performed, and information recording was performed. A high-density recording in the ordered molecular [system] was possible.



Chalcone

Playback of the recording was performed by reading out the change in absorption at a wavelength of around 380-420 nm associated with dimerization of the guest molecule. Subsequently, when ultraviolet rays were applied at a wavelength of 270 nm for 1 h, depolymerization took place, and the recording was erased.

Application Examples 2-12

Instead of the chalcone described in Application Example 1, compounds indicated by Nos. 31-38 were used as guest molecules, and similar results were obtained. (Application Examples 2-9)

Furthermore, when chalcone was used as the guest molecule, and compounds indicated by Nos. 50-52 were used as host molecules, recording, playback, and furthermore, erasure of the recording were made possible as in Application Example 1. (Application Examples 10-12)

Application Examples 13-24

An optical recording medium that utilizes the photodimerization reaction of the guest molecule (2)

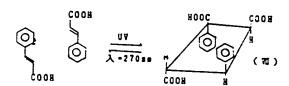
Total exposure was first performed for each optical recording medium described in Application Examples 1-12 using a high-pressure mercury lamp, and dimerization was performed for all of guest molecules. Ultraviolet rays with a wavelength of around 270 nm, which corresponds to the maximum absorption of the cyclobutane ring produced at the time of dimerization, were applied to the above-mentioned media to form a pattern, depolymerization of the guest molecule was carried out, and information recording was performed. Recording in the ordered molecular [system] was possible. Playback was accomplished by reading the change in absorption at wavelengths of approximately 380-420 nm associated with depolymerization of the guest molecule. It was confirmed that erasure of the recording, that

is, dimerization of all of the guest molecules, was possible by means of total exposure of the above-mentioned recording medium with a high-pressure mercury lamp.

Application Example 25

An optical recording medium that utilizes the photodimerization of the guest molecule (3)

Diacetylene diol indicated by No. 53 was used as the host molecule, and cinoamic acid was used as the guest molecule, and optical recording media with a recording layer comprised of a single complex molecular film, and built-in layers of 3, 5, 9, 15, and 19 layers were produced as in Application Example 1. Subsequently, total exposure was performed for above-mentioned films with a high-pressure mercury lamp, dimerization of the guest molecule (equation VII) was carried out, and polymerization of the host molecule (equation V) was carried out; subsequently, ultraviolet rays with a wavelength of 270 nm were applied according to a specified pattern, depolymerization of the guest molecule was performed, and information recording was accomplished.



Chalcone

Playback was accomplished by reading the change in absorption at wavelengths of approximately 380-420 nm associated with depolymerization of the guest molecule.

Subsequently, it was confirmed that erasure of the recording was possible by means of a total exposure of the above-mentioned recording medium with a high-pressure mercury lamp. Also, the recording medium that had been totally exposed with a high-pressure mercury lamp was dipped in alcohol for approximately 30 sec and recording/playback of information was performed as described above, and no problems were observed. In other words, it was confirmed that the chemical resistance of the recording medium can be increased when polymerization is performed for the host molecule.

Application Examples 26 and 27

Diacetylene diol indicated by No. 53 was used as the host molecule, and anthraldehyde was used as the guest molecule. They were dissolved in chloroform at a mole ratio of 1:2, and an optical recording media with recording layers comprised of a single complex molecular film, and built-in layers of 5, 9, 15, 31, 60, and 80 layers were produced as in Application Example 1.

Subsequently, X-ray radiation was performed of the optical recording medium produced, and the dimerization indicated by equation (IV) of the guest molecule was carried out, and information recording was performed. Recording in the ordered molecular [system] was possible. Playback was performed by reading the change in absorption at wavelengths of approximately 370-390 nm associated with dimerization of the guest molecule.

When ultraviolet rays with a wavelength of 313 nm were applied for 1 h, depolymerization took place, and it was confirmed that erasure of the recording was possible.

Similar results were obtained when the anthracene derivative indicated by No. 41 was used. (Application Example 27)

Table I				
۵				
	基本市场			
Mo. 43	No.10	n = 9 .	2 = 2	
N a . S 0	No.25	Z-COOH.	= 2	
No.51	No.15	m = 9 .	a = 2	
Na.52	No.30	z = соон .	2 = 4	
No.53	Na.7	m = 8 .	± = 8	
No.54	No. 4	a = 9 .	= 2	

Key: 1 Base skeleton

Brief explanation of the figures

Figures 1-3 show vertical cross section diagrams used for explanation of application examples of the recording medium of the present invention. Figure 1 shows the recording process, Figure 2 shows the playback process, and Figure 3 shows the erasure process, respectively. Figure 4 and Figure 5 are explanatory diagrams that shows the state of the inclusion complex of the present invention at the aqueous phase interface.

- 1...Host molecule
- 2...Guest molecule
- 3...Hydrophilic group
- 4...Long-chain alkyl group
- 5...Base
- 6...X-ray
- 7...Dimer
- 8...Visible rays
- 9...Ultraviolet rays
- 10, 11...Inclusion group
- 12, 13...Noninclusion group
- 14...Long-chain fatty acid group
- 15...Hydrophobic group
- 16...Aqueous phase

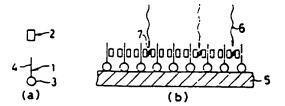


Figure 1

Figure 2

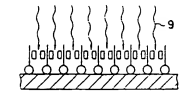


Figure 3

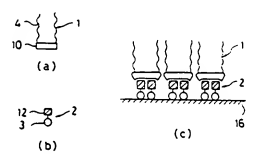


Figure 4

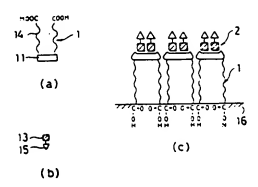


Figure 5

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②特 願 昭59-95575

②出 願 昭59(1984)5月15日

松 \blacksquare 宏 ⑫発 明 者 Ħ 宏 砂発 明 者 春 平 井 裕 砂発 明 者 ⑫発 明 者 西 村 生 健 明 者 ĭΤ ⑦発 明 中 桐 孝 志 砂発 老 キャノン株式会社 化出 頗 弁理士 若 林 忠 砂代 理

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朝 細

1. 発明の名称

記量媒体

2.特許請求の範囲

その分子内に親水性部位、硬水性部位及び包接部位を有するホスト分子と減ホスト分子に包接され光二量性を有するゲスト分子とからなる包接館体の単分子膜又は単分子層果積膜を担体上に形成して記録層とし、光により記録を行なうことを特徴とする記録媒体。

1. 是明の詳細な説明

(1) 技術分野

本発明は、包接遺体の単分子膜、乃至単分子層 環積膜の化学変化若しくは物理変化を利用して記 量を行なう記録媒体に関する。

(2) 背景技術

:

従来、有機化合物を記録層とする記録媒体としては種々のものが知られている。

別えば、有機化合物を薄膜にして記録層として 用いる光記録媒体については、例えば特開昭 56-16948号公報、特開昭58-125246号公報にも開示されている。いずれも有機色素を記録層とし、レーザビームにより記録再生を行なうレーチ記録媒体に関するものである。特に、特別密58-125246号公報に開示された媒体は、

一般式(I)

る。しかしながら、塗布法による場合、腰厚においては、300人程度が限界であり、腰内の分子分布、配向がランダムであることは解決しがたいことであった。

レジスト材料の一つとして光量子効率が大でかつ 受れた 展象力を有するものとして提案されていたジアセチレン化合物果積膜が、レジスト材料のみならず、 専課電気 - 光学デバイス、電気 - 音響デバイス、圧・無電デバイス等にも応用されることが、特別昭56-42229号公報、特別昭56-43220号公報などに示されている。

近時においては、ジアセチレン化合物累積膜のの製造方法の改良について特開昭58-111029 号公 選 公 記 で これている。かかる発明にて製造された照 を 別 が な を 財 重合 させて ジアセチレン 化合物累積膜は 無外線を 財 動 全 に より 重合 させて ジアセチレン 化合物 別 を 開 を 作り、 或はマスキングして 常外線を 関 形 を 節分 に 重合 させ、 未重合部分を 除去して 区 を 作り、 連 膜 光学 デバイス や 集積 回路 素子 と して 使 用 される。

た.

4

439

かからを解析し、1)各種の機能能性性能性を対象の欠点を解析し、2)その際、機能能性性能性的簡単に作製する方法、2)その際、は機能が、薄膜化した場合を提供が、3)上になくな関係を見いる方法、更には、3)上に、原構は存むに、特別な操作を行うことなり原構を発化して、特別な操作を行うことの原構を発明を対して、高度の記録と明かる成践を明かる成践を明かる成践を明かる成践を明かる成とを表明を表現して、高層像度の記録媒体を、容易にかつ高品質にという。

(3) 発明の開示

本発明の目的は、光により分子単位での化学変化若しくは物理変化を起こす様な高密度記録媒体を提供することにある。

また、この様な分子単位での高密度像形成を行うのに際して重要な因子となる媒体面内での分子配向に関して、従来例よりも秀遠な媒体を提供することにある。更には、上述記録媒体を製造する

しかし、これらはいずれもジアセチレン化合物に限るものであり、薄膜光学デバイスとして使用するときに、一度記録したものの消去の可能性については述べられていない。

一方、上述欠点を解決すべく、分子内に親末基、確永基及び少なくとも1個の不飽和結合を有する1種類の光重合性モノマーの単分子膜又は単分子層果種膜を基板上に形成して記録層としたことを特徴とする、反復使用可能な光記録媒体が特額昭58-190932 号の光記録媒体に示されている。

に当って、比較的簡単な操作変更により、 様々な 性質を有する媒体を提供することにある。

本発明の上記目的は、以下の本発明によって達成される。

その分子内に親水性部位、 積水性部位及び他分子との包接が可能な部位(包接部位)を有する分子 (ホスト分子に包接され、 光二量性を有する別種の分子 (ゲスト分子) とかか 担なる 包接 鎌体の単分子競又は単分子層 異 積 膜 を 足体上に形成して記録層とし、光により記録を行なうことを特徴とする記録媒体。

時間昭 60-239741(3)

種類の分子の内、ゲスト分子が、光により、二量化することが必要である。

ンジオール誘導体 (No.7~No.12. No.22 ~ No.27)、ハイドロキノン誘導体 (No.13 ~ No.15, No.28 ~ No.30) 等が利用し得るものとして挙げられる。 尚、以下の例における m, nは、正の整数を、こは、-CH3または-COOHを、Phは、-Ce Hsを示すものとする。

(ここで、X=H またはGBである。)

すなわち、分子内に親水性部位および硬水性部位を有するとは、例えば上式に於いてR1部及びR2部の何れか一方に親水性部位が存在し、他方に環水性部位が存在し、他方に現水性部位が存在するか、R1部及びR2部が両部以外の競りの部との関係に於いて共に親水性、若しくは環水性を示すことを言う。R1部及び、R2部のは環水性を示すことを言う。R1部及び、R2部のは環水性を示すことを言う。R1部及び、R2部のは環水性を示すことを言う。R1部及び、R2部のは環水性を示する場合には特に炭素原子数 1~30の脂肪酸が望ましい。

本発明に於けるホスト分子を更に具体的に示せば、例えば以下に列挙するアセチレンジオール誘導体 (No.1~No.8, No.18 ~No.21)、ジアセチレ

〔アセチレンジオール毎選供の刷〕

No. 1

No. 2

No. 3

$$CH_3-(CH_1)_m-O-\bigodot O+\bigodot O+O-(CH_1)_n-COOH$$

 $30 \ge m + n \ge 8$, $n \ge 1$

 $30 \ge m + n \ge 8$, $n \ge 1$

No. 5

T

Ø

 $30 \ge m + n \ge 8$, $n \ge 0$

No. 6

 $30 \ge m + n \ge 8$, $n \ge 0$

No. 7

$$30 \ge m + n \ge 5$$
, $n \ge 0$

No. 12

$$CH_3 - (CH_2)_m - 0 - \bigcirc - \bigcirc - C = C - C = C - C - C - \bigcirc - (CH_2)_m - COOH$$

 $30 \ge m + n \ge 5$, $n \ge 0$

No. 13

$$CH_3-(CH_3)_m$$
 OH OH

 $30 \ge m + n \ge 13$, $n \ge 0$

No. 14

$$CH^{1}-(CH^{4})^{m}-O$$
OH
OH
OH

 $30 \ge m + n \ge 9$, $n \ge 0$

No. 8

$$CH_{i}-(CH_{i})_{m}-Ph$$
 $C-C=C-C=C-C-(CH_{i})_{a}-COOH$
 OH
 OH

 $30 \ge m + n \ge 9$, $n \ge 0$

No. 9

$$CH_3 - (CH_2)_m - O - \bigcirc + \bigcap_{\substack{i \\ i \\ OH}} + C = C - C = C - C - O - (CH_1)_n - COOH$$

 $30 \ge m + n \ge 5, \quad n \ge 1$

No. 10

 $30 \ge m + n \ge 5$, $n \ge 1$

No. 11

 $30 \ge m + n \ge 9$, $n \ge 1$

No. 15

$$CH_3-(CH_2)_m-O-\bigcircO+\bigcircO+\bigcircO+$$

$$30 \ge m + n \ge 9$$
, $n \ge 0$

No. 16

30 ≥ n ≥ 5

No. 17

$$Z - (CH_t)_a - \overset{Ph}{\overset{\cdot}{C}} - C \equiv C - \overset{Ph}{\overset{\cdot}{C}} - (CH_t)_a - Z$$

$$\overset{\cdot}{OH} \overset{\cdot}{OH}$$

30 ≥ n ≥ 5

$$Z - (CH_t)_a - O - \bigcirc - \bigcirc - C - C - C - \bigcirc - O - (CH_t)_a - Z$$

30 ≥ n ≥ 4

No. 19

30 ≥ a ≥ 4

No. 20

30 ≥ n ≥ 4

No. 21

30 ≥ n ≥ 4

利用し得るゲスト分子の具体例としては、オレフィン化合物 (No.31 ~No.34)、ジオレフィン化合物 (No.35 ~No.38)、アントラセン誘導体 (No.33)、2-アミノビリジニウム (No.41)等が挙げられる。

以上挙げた化合物はポスト分子に長期アルギル 基や長期カルボン酸等を置換させて収水性や境水 性を導入した点を除けばそれ目体既知の化合物で あり、又、長期アルギル基等で修飾されていない ポスト分子が、種々のゲスト分子と結晶性の包 清体を形成する点に関しても、日本化学会造物、2 233頁-242頁(1983年)に述べられている。

ゲスト分子の光二量化反応を用いた記録媒体に

〔オレフィン化.合物の例〕

No. 31

$$\begin{array}{c}
X & O \\
CH = CH - C - R & (X = -H, -0 - CH, -P - CH, R = -H, -C, H, -OH, -OCH, -OC$$

No. 32

$$\begin{array}{c}
\bigcirc \\
\bigcirc \\
\bigcirc \\
\bigcirc \\
-CH = CH - C - R
\end{array}$$
(R = -H, -C₄H₁, -OH, -OCH₃)

No. 33

NC-CH=CH-CN

No. 34

 $R_1OOC-CH=CH-COOR_2$ ($R_1=R_2=CH_0$ \$ $\not\sim$ $R_1=CH_1$, $R_2=H$)

(ジオレフィン化合物の例)

No. 35

 A_{r}' - CH = CH - A_{r} - CH = CH - A_{r}'

時間昭 60-239741(6)

$$(Ar = \stackrel{N}{\longleftarrow} の場合は、Ar' = \stackrel{N}{\longleftarrow} , \stackrel{N}{\longleftarrow} ,$$
 $Ar = -\stackrel{N}{\longleftarrow} -$ の場合は、 $Ar' = \stackrel{N}{\longleftarrow} -$ 、 $Ar' = \stackrel{N}{\longleftarrow} -$

No. 36

No. 37

No. 38

$$R-CH=CH-C-CH=CH-R \qquad (R=C_0H_5-, -COOCH_5)$$

 $\Pi A = k T$

 (アントラセン請導体の例)

(アクリジニウム誘導体の例)

(2-アミノビリジニウム)

. 41

いて、本発明のゲスト分子を包接するホスト分子の単分子膜(これを単端体分子膜と呼ぶことにする)、若しくは単鏡体分子層累積膜の具体的な製法としては、例えば以下に示すA~Eの5法を挙げることができる。

 Z = - COON では末相上に形成される包接鎖体は、ホスト分子の現本性部位を末相に向けた第5 図に示すような状態で末相上に展開する。

単鏡体分子層を担体上に移すには、上述した垂っ 直浸せき法の他、水平付着法、回転円筒法などの 方法による。水平付着法は担体を水面に水平に接 触させて移しとる方法で、回転円筒法は、円筒型

の担体を水面上を回転させて単角体分子層を担体 表面に移しとる方法である。前述した垂直浸せき はでは、表面が親木性である担体を水面を横切る 方向に水中から引き上げるとホスト分子の提水基 が担体側に向いた単盤体分子層が担体上に形成さ れる。前述のように担体を上下させると、各行程 ごとに1枚ずつ単角体分子層が積み重なってい く。成職分子の向きが引上げ行程と提せき行程で 逆になるので、この方法によると各層間はポスト 分子の親木基と親木基、ホスト分子の硫木基と確 水基が向かい合うY型膜が形成される。それに対 し、水平付着法は、担体を水面に水平に接触させ て移しとる方法で、ホスト分子の韓木基が担体値 に向いた単盤体分子層が担体上に形成される。こ の方法では、累積しても、成膜分子の向きの交代 はなく全ての層において、疎水基が担体側に向い たX型膜が形成される。反対に全ての層において 親水芸が相体側に向いた黒薄膜は2型膜と呼ばれ

回転円筒法は、円筒型の担体を水面上を回転さ

せて単分子層を担体表面に移しとる方法である。 単分子層を担体上に移す方法は、これらに限定されるわけではなく、大面積担体を用いる時には、 担体ロールから水相中に担体を押し出していく方 法などもとり得る。また、前述した親水基、疎水 基の退体への向きは原則であり、退体の表面処理 等によって変えることもできる。

る。 又、 ゲスト分子の みが 機能性を持つ場合には、 このゲスト分子への化学的 体飾、 即ち、 頑水 基 や 親水 基 の導入を行わないので、 膜化に伴う 機能の 低下 は生じない。

- [B] 水溶性を示すゲスト分子を水相に溶解させる。次にホスト分子を溶剤に溶解せしめてこれを水相上に展開させる。この時间時にホスト分子・ゲスト分子間で包接鏡体形成が行われて腹状に折出する。ホスト分子とゲスト分子の組み合わせ及び以下の成膜操作については[A]に示した方法に選ずる。
- [C] 水容性を示すゲスト分子を水相に容解させる。次に、目的とする包接遺体のホスト分子とゲスト分子とを溶剤に溶解し、これを水相上に展開させて包接遺体を膜状に折出させる。ホスト分子とゲスト分子の組み合わせ及び以下の成膜操作については[A]に示した方法に準ずる。
- 【D】ホスト分子を溶剤に溶解しこれを水相中に 開きせる。その後、密閉系の装置を用いて気相 週、即ち装置内の空間をゲスト分子ガス雰囲気と

上述の方法によって担体上に形成される単角体 分子膜及び単角体分子層果發膜は高密度で記録を有しており、これらの膜で記録を で記録であることによって、包接角体の破綻に応気を 構成することによって、包接角体の破綻に応気の で光記録、熱的記録、電気的記録あるいは磁度を で光記録、高密度で高解像度の記録機能を する記録媒体を得ることができる。

み合わせ及び以下の成膜操作については、 【A】

に示した方法に準ずる。・

単角体分子層果發膜では、二量化物が容易に得られるのみならず、二量化反応に伴って生成が考しいません。 される各種の異性体差しくは構造体の唯一種としか 生成されない。即ち、包接鏡体層に於けるがスト 分子間の立体配列は、極めて整然としている。また、二量化した後は、暗所下でも解重合とは第二 だ、非照射部位は単量体のままであるので、記録が成 される。

記録された情報の読み取りは例えば可視光の照射によって行なう。すなわち、重合によって単量体時の共役系が崩れるので、可視光の吸収被長に変化をきたす。最大吸収被長は低波長側にシフトするので、吸収スペクトル変化を読みとることにより情報の再生が行なわれる(第2図)。

再生は、可視光による吸収スペクトル変化の読み取り以外にも、単量体時と二量化後の体限変化 又は屈折事変化をシュリーレン法により読みとることも可能である。この方法は、単量体時と二量化後の体験変化又は屈折事変化の大きい構造を有 作成した単雄体分子額及び単雄体分子層果積膜 を記録媒体の記録層として用いる場合以下に示す 様に記録が行われる。

例えばホスト分子に No. 7~15、 No. 22~30の何れかと、 No. 31-38 帝の光二量性二重結合を有するゲスト分子とを組み合わせるとホスト分子が デスト分子の組成比(mol 比)が1:2の包接路 体が形成される。この単循体分子膜、 若しく は、単端体分子層果發膜にあるパターンに従って ガンマ線、 X線、 紫外線など重合に必要な な エル ギーを供給しうる光を照射すると照射部位におい で 工式に示すようにゲスト分子間で二量化反応が おこる。

これらの反応は互いに崩壊する不望和結合の距離 が4 人以下のときおこり得るものであるが、 先に 述べた様な方法で作成された単錆体分子膜又は、

する化合物の単館体分子膜または単館体分子層製 強膜のときには特に適している。また、単館体分 子膜または単端体分子層製積膜を基板の上に単端体分子層製積膜を基板の上に を形成し、その上に単館体分子膜または単位体 分子層累積膜を形成することにより、単量体 分子層累積膜を形成することにより、単位体 分子層累積度の差を電気的に読み取ることも可能 である。

二量体はシクロプタン環に基づく、被長270mmの吸収を持つが、この被長270mmの無外光を無射することによって二量体は元の単量体二分子に戻る。従って一度記録された情報の除去が可能となる。(第3図)

なお光二量性のゲスト分子としてアントラセン 誘導体 No.39の如き化合物を用いる事もできる。 この場合光二量化反応は呼ばに従って進行する。

ス、ホスト分子として No.7を用いた場合には、X線、ガンマ線、無外線等の重合に必要なエネルギーを供給し得る光を無射すると、無射器位に於いてV式に示す様にホスト分子間で重合がおこり、ポリジアセチレンが形成される。

ものを使用する必要がある。使用することのできる担体の例としては、ガラス、アルミニウムなどの金属、プラスチック、セラミックなどが挙げられる。

担体上の単鏡体分子膜または単鏡体分子層累積 段は、十分に強く固定されており担体からの割 意、剥落を生じることはほとんどないが、 付着力 を強化する目的で担体と単鏡体分子膜または単鏡 体分子層累積膜の間に接着層を設けることもでき る。さらに単鏡体分子層形成条件例えば水相の水 素イオン濃度、イオン程、水温、担体上げ下げ速 度あるいは表面圧の選択等によって付着力を強化 することもできる。

単分子膜または単分子層累積膜の上に保護膜を 設けることは、単分子膜または単分子層累積膜の 化学的安定性を向上させるためには、好ましいこ とであるが、成膜分子の選択によって保護膜は設 けても設けなくてもよい。

以下に本発明の実施例を示して更に具体的に設明する。 No.49~ No.54の化合物は、第1表に示

を飛躍的に増大せしめることが可能である。 等に 耐寒品(耐容剤)性が増大する。かかる、全面露 光により、ゲスト分子が光二量性を示す場合には ゲスト分子も、二量化してしまうが、これを光に 足 選 体 と し て 用 い る 瞬 に は、前 述 例 と は 逆 に デ し ターンに 従ってシクロブタン環の 吸収 設 長 に 等 し い 波 長の 光 (紫外光)を 照射して 解重合 さ と により 記 録乃至表示が出来る 訳である。

以上述べた記録媒体に於いて顧厚は、特に 190~3000Åのものが好ましい。

これら成膜方法はその原理からも分る通り、非常に簡易な方法であり、上記のような優れた記録 機能を有する記録媒体を低コストで提供すること ができる。

以上述べた、本免明における単議体分子膜また は単菌体分子果硬膜を形成する担体は特に限定さ れないが、担体表面に界面活性物質が付着してい ると、単遠体分子層を水面から移しとる時に、単 遠体分子膜が乱れ良好な単鏡体分子膜または単鏡 体分子層累積膜ができないので担体表面が清浄な

す .

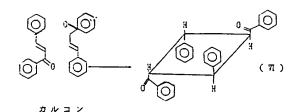
実施例 1

ゲスト分子の光二量化反応を用いた光記録媒体 (1)

ホスト分子として No.43のジアセチレオンジ オールと、ゲスト分子としてカルコンをモル比 1:2の割合でクロロホルムに溶かした後、 pH6.5 、 塩化カドミウム濃度 4 × 10 [★] N の水相上 に展開させた。溶媒のクロロホルムを蒸発除去 後、表面圧を35dynes/cmまで高めて、包接錯体を 避状に折出させた。この後、表面圧を一定に保ち ながら、表面が十分に清浄で製水性となっている ガラス基版を上下速度7ca/min にて水面を横切る 方向に静かに上下させ、単箇体分子膜を基板上に 移し取り、単盤体分子膜及び3,5,9,15,19 **港に果積した単蒲体分子層累積膜を記録層とする** 光記量媒体を製造した。この異稜行程に於いて基 版を永相から引きあげる郡废に、30分間以上放置 して、基板に付着している水分を蒸発除去した。 なお成蹊装置としては、英国 Joyce社製の

Laagmuir-Troughtを使用した。

作成した光記録媒体にパターンに従って、X級無財を行い、式町に示すゲスト分子の二量化反応を行ない情報を記録した。分子オーダーの高密度 記録が可能であった。



記録の再生はゲスト分子の二量化に伴う被長 380 nm~ 420 nm付近の吸収変化を読み取る事により行った。 次いで波長 270 nm の紫外光を 1 時間 照射したところ、解重合がおこり、記録が消去がされた。

実施例 2~12

実施例 1 のカルコンの代わりにゲスト分子として No.31~ No.38の化合物を用いた場合も何様の

岩果を得た。 (実施例 2~3)

ス、ゲスト分子をカルコンとし、ホスト分子をNo.50~ No.52とした場合にも、実施例 1 と何様、光に依る記録、再生、更には、記録の消去が可能であった。(実施例 10~12)

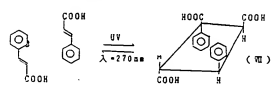
灵施例13~24

ゲスト分子の光二量化反応を用いた光記録媒体 (2)

実施例 1 ~12で述べた各光記録媒体を先づ高日田 水銀灯により全面露光せしめて、全てのゲストで 会工量化した。かかる媒体に二量化に伴っる。 ではされたシクロプタンでの吸収機大に無射して、 と270mm 付近の無外光を収収した。 分子を解重合し情報を記録の再生は近の再生は近の再生は近の再生は近の再生は近の再生は近のの配録が可能であった。 更にすることを 取変化を読み取る事により行った。 更に 水銀灯を用いて上記録媒体を全面電光トトの 取変化を読み取る事により行った。 更に 水銀灯を用いて上記録媒体を全面電光トトース 取変化を読み取る事により行った。 更に 水銀灯を用いて上記録媒体を全面電光トトース 最近により、記録を をことを確認した。

実施例 25

ゲスト分子の光二量化反応を用いた光記録媒体 (3)



定量の再生は、二量化したゲスト分子の原重合 に伴う波長380~420 □■付近の吸収変化を提み取る事により行った。 更に再び高圧水銀灯を用いて上記録媒体を全面 選光することにより記録を商去することが可能で あることを確認した。なお一度高圧水銀灯で全面 選光した本記録媒体を、アルコール中に約30秒間 浸漬した後、上記方法により情報の記録/再生を 行ったが、特に問題点はなかった。即ち、ホスト 分子を重合させる事により該記録媒体の化学的強 度が大となることが確認された。

実施例28.27

ホスト分子として No.53のジアセチレンジォール、ゲスト分子としてアントラアルデヒドをモル比1:2の割合でクロロホルムに容かし、実施例1と同様に操作により単分分子膜及びこれを5,9,15,31,80,80層に累積した単分分子層累積膜を記録層とする光記量媒体を製造した。

作成した光記量媒体にパターンに従ってX線照射を行ない、式(IT)に示すゲスト分子の二量化反応を行ない情報を記録した。分子オーダーの記録が可能であった。記録の再生はゲスト分子の二量化に伴う彼長370~ 390mm付近の吸収変化を禁

み取る事により行った。次いで、波長 J1Jamの 紫 外光を 1 時間照射したところ解重合が起こり、記 録の消去が可能であることを、確かめた。

No.(1のアントラセン誘導体を用いた場合にも 同様の結果を得た。(実施例27)

羽 1 表

	基本骨格	
No.49	Na.10	m = 9 , n = 2
No.50	No. 25	Z • C O O H . n = 2
No.51	Na.15	m = 9 . n = 2
No.52	No.30	Z=COOH, n = 4
No.53	No. 7	m = 8 , n = 8
Но.54	Na. 4	m = 9 . n = 2

4.図面の簡単な説明

第1図~第3図は、本発明に係る療託誘揮体の 実施例を説明する厳断面図であり、各々、第1図 は魔脈城 多記録》過程、第2図は再生過程、第3 図は消去過程を示しており、第4図~第5図は本 発明に係る包接着体の水相上に於ける状態を説明 する説明図である。

1 --- ホスト分子

2 --- ゲスト分子

3 --- 親永性部位

4 --- 長額アルキル部位

5 --- 基板

6 --- X 🖳

7 --- 二量化部位

8 --- 可视光

16

10. 11--- 包接部位

12, 13--- 被包接部位

14--- 長額脂肪酸部位

15--- 蘋水性部位

16--- 水相

